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(9) Organic nonlinear optical substance.

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Description

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1. Field of the Invention

The present invention relates to a novel nonlinear optical material naving an increased second harmonic generating ability and usable for an optical switch, an optical memory in an optical data information processing or optical communication system, or an optical bistable element to be used in optical signal operation. Further, it pertains to a novel nonlinear optical crystalline material containing a salt obtained by the reaction of a conjugated aromatic carboxylic acid compound with an optically active amine.

2. Description of the Related Art

The nonlinear optical effect refers to, for example, a secondary or higher effect of the magnitude of an applied electric field exhibited by the relationship of an electric polarizing response of a substance, which is only primarily proportional to the magnitude of the applied electric field, when, for example, a strong photoelectric field such as a laser beam is applied to the substance.

A secondary nonlinear optical effect may include a second harmonic generation of converting the wavelength of an incident light to a 1/2 wavelength, a parametric oscillation which converts a light with one kind of wavelength into a light with two kinds of wavelengths, and a secondary light mixing which on the contrary generates a light with one kind of wavelength from a light with two kinds of wavelengths. Due to these various characteristics, materials having a nonlinear optical effect will be for use as an optical switch, optical memory in optical data/information processing, or such elements as optical bistable element, optical switch, etc. to be used in optical signal operations.

Generally speaking, in this field of the art, inorganic materials, primarily LiNbO₂, have been studied and investigated, but inorganic materials had a drawback such that great difficulty is encountered in forming a desired optical element, because of such shortcomings as their performance indices which are not so great, small response speed, no good form workability, great hygroscopicity, low stability, etc.

In recent years, in contrast to these inorganic materials, researchers are becoming more interested in application of organic materials. This is because the response of organic materials is based primarily on the π electron polarization, whereby the nonlinear effect is great, and also the response speed is great, as confirmed and reported in the art. For example, a large number of studies are reported in the ACS Symposium Series Vol. 233, 1983. The secondary nonlinear optical characteristics to be dealt with in the present invention, which is a third rank tensor, cannot be evoked if a symmetric center exists in the molecular or the crystal. For this reason, in the case of organic materials, even when they may have a structure exhibiting an excellent nonlinear effect at molecular level, they must be formed into crystals or solid state for using the second harmonic generation as the practical mode. However, at such stage of solidification, an inversion symmetrical structure will be frequently formed preferentially, whereby there has been involved the problem that a nonlinear optical effect as an optical element can not be exhibited.

40 SUMMARY OF THE INVENTION

having the formula (1):

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Accordingly, an object of the present invention is to solve the above-mentioned problems of the prior art and to provide an organic crystalline compound for various nonlinear optical elements, and having an increased second harmonic generating ability, a high molecular polarizing ability, and having no inversion symmetry.

Another object of the present invention is to provide an organic crystalline compound having excellent second harmonic generating ability required for a material for forming various optical signal processing elements.

Other objects and advantages of the present invention will be apparent from the following description. In accordance with the present invention, there is provided an organic nonlinear optical substance

$$A \leftarrow CR^{1} = CH \rightarrow \frac{1}{n} CH = C - C - B \qquad (I)$$

wherein

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R' represents -H or -CH₅, n is 0, 1, or 2. A represents Z'-Ar.

$$z^{2}$$
 z^{3}
 z^{4}
Ar-, z^{2} , or (z^{5})
 z^{1}
 z^{2}

wherein Ar represents a 6 - 14 membered aromatic group including a heterocyclic ring or bisphenylene type ring. Z' represents H-, $R^{\xi}R^{\xi}N$ -, $R^{2}O$ -, $R^{\xi}S$ -, NC-, $R^{\theta}OCO$ -, $R^{10}COO$ -, $O_{\xi}N$ -, $R^{11}R^{12}NOC$ -, $R^{13}CO$ -, R^{1

B represents -OH+Amine where Amine represents an optically active amine; -NR⁴Y where R⁴ represents -H or a single bond; Y represents -(CH₂) $_p$ CQ¹Q²Q³ where p is 0 or 1; Q⁷, Q², and Q³ are different and represent -H, a C₁-C₅ alkyl, phenyl, naphthyl, -OH, -CH₂OH, -COOR²⁵, -CNR²⁶R²⁷, a residue of an α -amino acid from which an amino group is removed, where R²⁵ to R²⁷ independently represent -H or -C·-C₈ hydrocarbon residue, or Y represents -CQ⁴Q⁵Q⁶ where Q⁴, Q⁵ and Q⁶ are different and Q⁴ and Q⁵ are as defined for Q¹, Q² and Q³ and Q⁶ represents

$$-(CH_2)_{\overline{q}}$$

of which one bond is linked to the bond of R4 where q is an integer of 1 to 4.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

Generally speaking, the second harmonic generating ability is greater in a longer conjugated system, in which polarization within the molecular is greater and also contribution of the polarization is greater, but with elongation of the conjugated length, the absorption maximum is shifted toward the longer wavelength side, whereby correspondence to a 1/2 wavelength of the incident light may occur. In this case, optical damage may occur which absorbs the second harmonic generated and changes the refractive index, chemical denaturation, or combustion by an absorption of heat energy. Accordingly, it is frequently disadvantageous to simply elongate the conjugated length. For example, a compound increased in molecular polarization due to an insertion of a carboxyl group represented by the formula (I) shown below, a group having a high electron attractability such as a cyano group, and further various substituents into benzene nucleus may be expected to have large nonlinearity as the result of shifting effect of electron arrangement within the ring. but practically has a structure with an inversion asymmetry center due to a large molecular polarization, whereby frequently a generation of the second harmonic can not be observed. Generally speaking, it is a difficult technique to control the crystalline structure, particularly to form a crystal form which will collapse the symmetry center. Accordingly, while an excellent nonlinear susceptibility may be expected to be 50 possessed at molecular level, most examples prove to be no longer effective as the second harmonic generating material. As the result of intensive studies, as shown in the present invention, by use of an optically active amine as the basic substance to introduce its optically active asymmetric structure thereof as the carboxylic acid salt, and consequently a structure without an inversion asymmetry center can be prepared to accomplish the present invention. As a consequence, the great nonlinear susceptibility at molecular level can be exhibited as such as the crystal structure, and this may be considered to contribute greatly to applications in this field of the art. ျမှား မြောင်းကောင်း သည်။ သည် သည် သည် သည် သည် သည် သည် သည်။

the same carbon atom. Also for the molecular polarizations to interfere with each other, it is desirable that a conjugated system exists, but an elongation of the conjugated length will result in an elongation of the absorption maximum to the longer wavelength side, whereby there is the possibility that the damage may be formed due to the incident light wavelength or the second harmonic wavelength. For this reason, the conjugated length should not be too long.

From the above-mentioned standpoints, it has been found that α -cyanoacrylic acid derivatives having a conjugated group are effective as an acid moiety having a conjugated system. Furthermore, the compounds having the formula (I) including the optically active amine satts or amides are effective for providing asymmetry center, as mentioned above

According to a first embodiment of the present invention, the organic nonlinear optical substance has the general formula (I) wherein R' represents H, Z' represents H, R' R'N-, R'O-, R'S-, NC-, R'S-COO-, Ω -N-, R'COO-, R''R'OCO-, R''S-CO(R'')N-, or R'S-, Ar represents an aromatic group having 5 to 14 carbon atoms, and B represents -OH+Amine* wherein Amine* represents an optically active amine selected from the group consisting of 1-phenylethylamine 1-(α -naphthyl)ethylamine, 1-phenyl-2-methylethylamine, 1-phenyl-2-aminopropane, and brucine.

These amines are strong bases and, therefore, easily react with carboxylic acids to form stable salts in any conventional method.

The optically active amines may be dextrorotatory or levorotatory. Formation of the salt may be carried out by conventional neutralization reaction, which may be in either state of a solution or a solid phase.

For maintaining the purity of the optical activity, it is not preferable to carry out the reaction at too high a temperature, but it is desirable to contrive to inhibit heat generation during salt formation. Salt formation will frequently give a product which differs in solubility to a great extent from the starting material, and therefore existence of salt formation can be easily confirmed and its purification can be also easily done.

The optically active amine salt of the carboxylic acid thus obtained assumes a crystalline state, having excellent moldability, and can be formed into various elements under the crystalline form as such or as a solid solution, which can be then applied to the nonlinear optical application field.

The organic nonlinear optical substances according to the first embodiment of the present invention have an extremely large second harmonic ability. The carboxylic acid moiety of this compound can be preferably derived from the caboxylic acids having the structure (II)

$$z^{1} - A_{r} - (CH = CH) - CH = C - COOH (II)$$

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Examples of such carboxylic acid may include: substituted phenyl 2-cyanopropenoic acid derivatives such as 3-phenyl-2-cyanopropenoic acid, 3-(p-dimethylaminophenyl)-2-cyanopropenoic acid, 3-(p-aminophenyl)-2-cyanopropenoic acid, 3-(p-diethylaminophenyl)-2-cyanopropenoic acid, 3-(p-dipethylaminophenyl)-2-cyanopropenoic acid, 3-(p-monomethylaminophenyl)-2-cyanopropenoic acid, 3-(p-monomethylaminophenyl)-2-cyanopropenoic acid, and the m- or o- substituted derivatives;

3-(p-methyloxyphenyl)-2-cyanopropenoic acid, 3-(p-ethyloxyphenyl)-2-cyanopropenoic acid, 3-(p-propyloxyphenyl)-2-cyanopropenoic acid, 3-(p-butyloxyphenyl)-2-cyanopropenoic acid, 3-(p-pentyloxyphenyl)-2-cyanopropenoic acid, 3-(p-decanoxyphenyl)-2-cyanopropenoic acid, and the m-or ε- substituted derivatives;

3-(p-methylthiophenyl)-2-cyanopropenoic acid, 3-(p-ethylthiophenyl)-2-cyanopropenoic acid, 3-(p-propyl-thiophenyl)-2-cyanopropenoic acid, 3-(p-butylthiophenyl)-2-cyanopropenoic acid, 3-(p-pentylthiophenyl)-2-cyanopropenoic acid, 3-(p-decanethiophenyl)-2-cyanopropenoic acid, and the m-or c- substituted derivatives,

3-(p-cyanophenyl)-2-cyanopropenoic acid, 3-(m-cyanophenyl)-2-cyanopropenoic acid, 3-(o-cyanophenyl)-2-cyanopropenoic acid, 3-(p-ethyloxycarbonylphenyl)-2-cyanopropenoic acid, 3-(p-ethyloxycarbonylphenyl)-2-cyanopropenoic acid, 3-(p-propyloxycarbonylphenyl)-2-cyanopropenoic acid, and the more of substituted derivatives;

3-(p-acetyloxyphenyli-2-cyanopropenoic acid, 3-(p-propionyloxyphenyli-2-cyanopropenoic acid, 3-(p-butanoylphenyli-2-cyanopropenoic acid and the m-or c- substituted derivatives.

3-(p-nitrophenyl)-2-cyanopropenoic acid. 3-(m-nitrophenyl)-2-cyanopropenoic acid. 3-(o-nitrophenyl)-2-

cyanopropenoic acid, and the m- or o- substituted derivatives.

3-(p-acetylaminophenyl)-2-cyanopropenoic acid 3-(p-propionylaminophenyl)-2-cyanopropenoic acid, and the m- or c- substituted derivatives:

3-(p-methylphenyl)-2-cyanopropenoic acid 3-(p-ethylphenyl)-2-cyanopropenoic acid 3-(p-propyl-ethylphenyl)-2-cyanopropenoic acid 3-(p-pentylphenyl)-2-cyanopropenoic acid 3-(p-n-hexylphenyl)-2-cyanopropenoic acid 3-(p-decanephenyl)-2-cyanopropenoic acid and the m- or o- substituted derivatives.

Substituted phenyl 2-cyano-2 4-pentadienoic acid derivatives such as

2-cyano-5-phenyl-2.4-pentadienoic acid. 2-cyano-5-(p-dimethylaminophenyl)-2.4-pentadienoic acid. 2-cyano-5-(p-diethylaminophenyl)-2.4-pentadienoic acid. 2-cyano-5-(p-dipropylaminophenyl)-2.4-pentadienoic acid. 2-cyano-5-(p-dibutylaminophenyl)-2.4-pentadienoic acid. 2-cyano-5-(p-monomethylaminophenyl)-2.4-pentadienoic acid. acid. 2-cyano-5-(p-monomethylaminophenyl)-2.4-pentadienoic acid. acid. and the m- or o- substituted derivatives:

2-cyano-5-(p-methyloxyphenyl)-2.4-pentadienoic acid.
 2-cyano-5-(p-ethyloxyphenyl)-2.4-pentadienoic acid.
 2-cyano-5-(p-butyloxyphenyl)-2,4-pentadienoic acid.
 2-cyano-5-(p-butyloxyphenyl)-2,4-pentadienoic acid.
 acid. and m-, o- substituted derivatives thereof;

2-cyano-5-(p-methylthiophenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-ethylthiophenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-butylthiophenyl)-2,4-pentadienoic acid, and m-, o- substituted derivatives thereof;

5-(p-cyanophenyl)-2,4-pentadienoic acid, and m-, o- substituted derivatives thereof;

2-cyano-5-(p-methyloxycarbonylphenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-ethyloxycarbonylphenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-propyloxycarbonylphenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-butyloxycarbonylphenyl)-2,4-pentadienoic acid, and m-, o- substituted derivatives thereof;

2-cyano-5-(p-acetyloxyphenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-propionyloxyphenyl)-2,4-pentadienoic acid, and m-, o- substituted derivatives thereof:

2-cyano-5-(p-nitrophenyl)-2.4-pentadienoic acid, and m-, o- substituted derivatives thereof;

2-cyano-5-(p-dimethylamidophenyl)-2.4-pentadienoic acid, 2-cyano-5-(p-diethylamidophenyl)-2.4-pentadienoic acid,

2-cyano-5-(p-dipropylamidophenyl)-2,4-pentadienoic acid. 2-cyano-5-(p-dibutylamidophenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-amidophenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-amidophenyl)-2,4-pentadienoic acid, and m-, o- substituted derivatives thereof;

2-cyano-5-(p-acetylaminophenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-propionylaminophenyl)-2,4-pentadienoic acid and m-, o- substituted derivatives thereof;

2-cyano-5-(p-methylphenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-ethylphenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-propylphenyl)-2,4-pentadienoic acid, 2-cyano-5-(p-butylphenyl)-2,4-pentadienoic acid and m-, o-substituted derivatives thereof;

Substituted phenyl 2-cyanc-2,4,6-heptatrienoic acid derivatives such as:

2-cyano-7-phenyl-2,4,6-heptatrienoic acid, 2-cyano-7-(p-dimethylaminophenyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(p-diethylaminophenyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(p-dipropylaminophenyl)-2,4,6-heptatrienoic acid.

2-cyano-7-(p-dibutylaminophenyl)-2,4, 6-neptatrienoic acid, 2-cyano-7-(p-aminophenyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(p-monomethylaminophenyl)-2,4,6-heptatrienoic acid, and m-, o- substituted derivatives thereof:

2-cyano-7-(p-methyloxyphenyl)-2.4.6-heptatrienoic acid. 2-cyanc-7-(p-ethyloxyphenyl)-2.4.6-heptatrienoic acid. 2-cyanc-7-(p-butyloxyphenyl)-2.4.6-heptatrienoic acid. 2-cyanc-7-(p-butyloxyphenyl)-2.4.6-heptatrienoic acid. and m- (- substituted derivatives thereof;

2-cyano-7-(p-methylthiophenyl)-2,4,6-heptatrienoid acid, 2-cyano-7-(p-ethylthiophenyl)-2,4,6-heptatrienoid acid.

5c 2-cyano-7-(p-propylthiophenyl)-2.4.6-heptatrienoic acid, 2-cyano-7-(p-butylthiophenyl)-2.4.6-heptatrienoic acid and m-, o- substituted derivatives thereof,

2-cyano-7-(p-cyanophenyl)-2.4,6-heptatrienoic acid. and m-, c- substituted derivatives thereof,

2-cyano-7-(p-methyloxycarbonylphenyl)-2,4,6-heptatrienoic acid.

2-cyano-7-(p-ethyloxycarbonylphenyl)-2.4.6-neptatrienoic acid. 2-cyano-7-(p-propyloxycarbonylphenyl)-2.4.6-heptatrienoic acid. 2-cyano-7-(p-biityloxycarbonylphenyl-2.4.6-heptatrienoic acid. and m- or o- substituted derivatives

Examples of such carboxylic acid include 3-(3.4-dimethoxyphenyli-2-cyanopropenoic acid. 3-(3.4-diethoxyphenyl)-2-cyanopropenoic acid, 3-(3,4-dipropyloxyphenyl)-1-cyanopropenoic acid, 3-(2,4-dimethoxyphenyl)-2-cyanopropenoic acid 3-(2 4-diethoxyphenyl)-2-cyanopropenoic acid, 3-(2.4-dipropyloxyphenyl)-2-cyanopropenoic acid, 3-(3.4-dimethylthiophenyl)-2-cyanopropenoic acid, 3-(3,4-diethylthiophenyl)2cyanopropenoic acid. 3-(3,4-dipropylthiophenyl)-2-cyanopropenoic acid. 3-(2,4-dimethylthiophenyl)-2cyanopropenoic acid. 3-(2.4-diethoxyphenyl)-2-cyanopropenoic acid. 3-(2.4-dipropylthiophenyl)-2cyanopropenoic acid. 3-(3,4-dimethylaminophenyl)-2-cyanopropenoic acid. 3-(3,4-diethylaminophenyl)-2cyano-1-propenoic acid, 3-(3.4-dipropylaminophenyl)-2-cyanopropenoic acid, 3-(2.4-dimethylphenyl)-2cyanopropenoic acid, 3-(2,4-diethylaminophenyl)-2-cyanopropenoic acid, 3-(2,4-dipropylaminophenyl)-2cyanopropenoic acid, 3-(3,4-dinitrophenyl)-2-cyanopropenoic acid, 3-(2,4-dinitrophenyl)-2-cyanopropenoic acid, 5-(3,4-dimethoxyphenyl)-2-cyano-2,4-pentadienoic acid, 5-(3,4-diethoxyphenyl)-2-cyano-2,4-pentadienoic acid, 5-(3,4-dipropyloxyphenyl)-2-cyano-2,4-pentadienoic acid, 5-(2,4-(dimethylthiophenyl)-2cyano-2,4-pentadienoic acid, 5-(2,4-diethylthiophenyl)-2-cyano-2,4-pentadienoic acid, 5-(2,4-dipropylthiophenyl)-2-cyano-2,4-pentadienoic acid, 5-(3,4-dimethylthiophenyl)-2-cyano-2,4-pentadienoic acid, 5-(3,4-dimethylthiophenyl)-2-cyano-2,4-dimethylthiophenyllhio diethylthiophenyl)-2-cyano-2,4-pentadienoic acid, 5-(3,4-dipropylthiophenyl)-2-cyano-2,4-pentadienoic acid, 5-(2.4-dimethylthiophenyl)-2-cyano-2,4-pentadienoic acid, 5-(2,4-diethylthiophenyl)-2-cyano-2,4-pentadienoic 5-(2,4-dipropylthiophenyl)-2-cyano-2,4-pentadienoic acid, 5-(3,4-dinitrophenyl)-2-cyano-2,4-pentadienoic acid, 5-(2,4-dinitrophenyl)-2-cyano-2,4-pentadienoic acid, 7-(3,4-dimethoxyphenyl)-2-cyano-2,4,6hexatrienoic acid, 7-(3.4-diethoxyphenyl)-2-cyano-2,4,6- hexatrienoic acid, 7-(3,4-dipropyloxyphenyl)-2cyano-2,4,6-hexatrienoic acid, 7-(2,4-dimethylthiophenyl)-2-cyano-2,4,6-hexatrienoic acid, 7-(2,4-diethylthiophenyl)-2-cyano-2,4,6-hexatrienoic acid, 7-(2,4-dipropylthiophenyl)-2-cyano-2,4,6-hexatrienoic acid, 7-(3,4-dimethylthiophenyl)-2-cyano-2,4,6-hexatrienoic acid, 7-(3,4-diethylthiophenyl)-2-cyano-2,4,6-hexatrienoic acid, 6-(3,4-dipropylthiophenyl)-2-cyano-2,4,6-hexatrienoic acid, 6-(3,4-dinitrophenyl)-2-cyano-2,4,6-hexatrienoic acid, 7-(2,4-dinitrophenyl)-2-cyano-2,4.6-hexatrienoic acid and the like. The mutual position of the double bonds of these conjugated carboxylic acids should be preferably trans-form, to obtain a stable structure and also for exhibiting the nonlinear optical effect but is not limited thereto.

The organic substance of the present invention is obtained as a solid with a crystalline structure not having an inversion symmetrical center according to the constitution as described above, and an excellent nonlinear susceptibility at the molecular level is exhibited as such also in the crystalline structure, thus exhibiting an excellent nonlinear optical effect such as a generation of the second harmonic at a high level, and therefore, this substance can be utilized for forming an optical signal processing element.

According to a fourth embodiment of the present invention, the organic nonlinear optical substance has the general formula (I) wherein R' represents H, A represents

$$z^{2}$$
 z^{3}
 \rightarrow Ar
 z^{4}

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wherein one of Z^1 , Z^2 and Z^4 represents H- and trie remainder independently represents $C_1 - C_2 - C_3 - C_4 - C_4 - C_4 - C_5 - C_5$

The optically active amine salt of the carboxylic acid thus obtained assumes a crystalline state, having excellent moldability, and can be formed into various elements under the crystalline form as such or as a solid solution, which can be then applied to the nonlinear optical application field. The organic nonlinear optical substances according to the first embodiment of the present invention have an extremely large

The carboxylic acid moiety of this compound can be preferably derived from the carboxylic acids having the structure (III)

According to a fifth embodiment of the present invention, the organic nonlinear optical substance has the general formula (I) wherein R° represents H, A represents Z°Ar-, Z° represents H- R°R°N-, R°O-, R°O-, R°O-, R°O-, R°COO, R°R°NOC-, R°R°

The moiety -NR⁴Y includes, for example, α -chiral substituted primary alkylamines (i.e., p=0) having 4 to 20 carbon atoms such as 1-methylpropylamine, 1-ethylpropylamine, 1-methylbutylamine, 1-methylpentylamine, 1-phenylethylamine, 1-(α -naphthyl)ethylamine, and 1-(hydroxymethyl)propylamine, β -chiral substituted primary alkylamines (i.e., p=1) having 4 to 20 carbon atoms such as 2-(methyl)butylamine, 2-methylpentylamine, 2-phenylpropylamine, 2-(α -naphthyl)propylamine, and 2-(hydroxybutyl)amine prolinole, amino acid derivatives having 3 to 20 carbon atoms such as alkyl esters (e.g., methyl esters and ethyl esters) of α -amino acids, α -amino acid amides, α -amino acid anildes, or peptides derived from the same or different α -amino acids. It should be noted, however, that the presence of a carboxyl group as an acid is not preferable from the reaction standpoints. Examples of the typical optically active (either dextrorotatory or levorotatory) amino acids, usable in the present invention are alanine, leucine, isoleucine, ethionine, cysteine, serine, tyrosine, tryptophan, threonine, norvaline, norieucine, valine, histidine, phenylalanine, α -phenylglycine, methionine, and proline; monoamino dicarboxylic acids and carboamides such as aspartic acid, glutamic acid, asparagine, and glutamine; and diaminomonocarboxylic acids such as arginine, lysine, ornithine, canavanine, and hydroxylysine

The carboxylic acid moiety of this compound can be preferably derived from the carboxylic acids having the structure (II).

According to further embodiments of the present invention, the organic nonlinear optical substances have the following general formula (I):

(i) R' represents H-, A represents Z'-Ar-, Z' represents RSR6N-, RZO-, R8S-, NC-, RSOCO-, R19COO-, R11RCON-, R13CO(R14)N-, or R15, Ar represents an aromatic group having 6 to 14 carbon atoms, and B represents -OH+Amine* wherein Amine* represents an optically active α -amino acid or the derivative thereof, which can include those as mentioned above

(ii) R' represents H, A represents

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$$z^{2}$$
 z^{3}
Ar

where one of \mathbb{Z}^2 , \mathbb{Z}^3 , and \mathbb{Z}^4 represents H or substituted C $-C_E$ alkyl, the remainder of \mathbb{Z}^2 , \mathbb{Z}^3 and \mathbb{Z}^4 represents together methylene dioxy group wherein the dioxy groups are bonded to the adjacent positions of Ar, Ar represents an aromatic group having 6 to 14 carbon atoms, and B represents $-OH+Amine^*$ wherein Amine* represents an optically active amine selected from the group consisting of 1-phenylethylamine, 1- α -naphthyl)ethylamine, 1-phenyl-2-methylethylamine, 1-phenyl-2-aminopropane, brucine, 1-butanol, 1-amino-2-propanol, 2-amino-1-propanol, 2-amino-1-(p-nitrophenyl)-1,3-propanediol, 2-dimethylamino-1-phenyl-1-benzyl-1-propanol, 1-(N,N-dimethylamino)-1-phenyl-propylamine, and a residue of an optically active α -amino acid and the derivative thereof, or the above-mentioned NR⁴ Y.

The carboxylic acids forming the organic nonlinear optical substances of this embodiment can be represented by the formula (IV)

$$z^{2}$$
 $O - Ar + CH = CH + CH = C - COOH$
 $CH_{2} - O$
 $CH_{2} - O$
 $CH_{3} - O$
 $CH_{4} - O$
 $CH_{4} - O$
 $CH_{5} - O$
 $CH_{5} - O$

Examples of such cartioxylic acids are 3-(3.4-dioxymethylenephenyli-2-cyanopropenoic acid. 2-cyano-5-(3.4-dioxymethylenephenyl)-2,4-pentadienoic acid. 2-cyano-7-(3.4-dioxymethylenephenyl)-2,4.6-

neptatrienoic acid, and piperonoyl derivatives.

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r,r

(iii) R1 represents H or CH₂. A represents R1 where R1 represents H or an alkyl group having 1 to 12 carbon atoms, and B represents -OH-Amine* wherein Amine* represents an optically active amine selected from the group consisting of 2-amino-1-butanol 1-amino-2-propanol, 2-amino-1-propanol, 2-amino-1-propanol, 2-dimetrylamino-1-phenyl-1-benzyl-1-propanol, and 1-(N.N-dimetrylamino)-1-phenyl-propylamine.

The carboxylic acids forming the organic nonlinear optical substances of this embodiment can be represented by the formula (V).

$$H \leftarrow CH_2 \rightarrow CR' = CH \rightarrow CH = C - COOH$$
 (V)

Examples of such carboxylic acid may include 3-alkyl substituted 2-cyano propionic acid derivatives such as 3-methyl-2-cyanopropenoic acid. 3-(n-propyl)-2-cyanopropenoic acid. 3-(n-butyl)-2-cyanopropenoic acid. 3-(n-hexyl)-2-cyanopropenoic acid;

5-Alkyl substituted 2,4-pentadienoic acid derivatives such as 2-cyano-5-methyl-2,4-pentadienoic acid, 2-cyano-5-(n-propyi)-2,4-pentadienoic acid, 2-cyano-5-(n-butyl)-2,4-pentadienoic acid, 2-cyano-5-(n-hexyl)-2,4-pentadienoic acid, 2-cyano-5-(n-hexyl)-2,4-pentadienoic acid, 2-cyano-5-(n-heptyl)-2,4-pentadienoic acid;

5-Alkyl substituted 2,4-pentadienoic acid derivatives such as 2-cyano-5-methyl-2,4-hexadienoic acid, 2-cyano-5-(n-propyl)-2,4-hexadienoic acid, 2-cyano-5-(n-butyl)-2,4-hexadienoic acid, 2-cyano-5-(n-pentyl)-2,4-hexadienoic acid, 2-cyano-5-(n-heptyl)-2,4-hexadienoic acid, 2-cyano-5-(n-heptyl)-2,4-hexadienoic acid:

7-Alkyl substituted 2,4,6-heptatrienoic acid derivatives such as 2-cyano-7-methyl-2,4,6-heptatrienoic acid, 2-cyano-7-(n-propyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(n-butyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(n-pentyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(n-hexyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(n-heptyl)-2,4,6-heptatrienoic acid;

5-Methyl 7-alkyl substituted 2,4,6-heptatrienoic acid derivatives such as 2-cyano-5-methyl-7-methyl-2,4,6-heptatrienoic acid, 2-cyano-5-methyl-7-(n-propyl)-2,4,6-heptatrienoic acid, 2-cyano-5-methyl-7-(n-pentyl)-2,4,6-heptatrienoic acid, 2-cyano-5-methyl-7-(n-hexyl)-2,4,6-heptatrienoic acid, 2-cyano-5-methyl-7-(n-hexyl)-2,4,6-heptatrienoic acid.

7-Alkyl substituted 2,4,6-octatrienoic acid derivatives such as 2-cyano-7-methyl-2,4,6-octatrienoic acid, 2-cyano-7-(n-propyl)-2,4,6-octatrienoic acid, 2-cyano-7-(n-butyl)-2,4,6-octatrienoic acid, 2-cyano-7-(n-hexyl)-2,4,6-octatrienoic acid, 2-cyano-7-(n-hexyl)-2,4,6-octatrienoic acid, 2-cyano-7-(n-heptyl)-2,4,6-octatrienoic acid.

(iv) R' represents H, A represents

$$(z^5) = \begin{bmatrix} c & -c \\ r & & \\ c & c \\ x & &$$

and B represents -OH+Amine* wherein Amine* represents an optically active amine selected from the group consisting of 1-phenylethylamine, 1-(a-naphthyliethylamine, 1-phenyl-2-methylethylamine, 1-phenyl-2-aminopropane brucine 2-amino-1-butanol 1-amino-2-propanol 2-amino-1-propanol 2-amino-1-(p-nitrophenyl)-1,3-propanediol, 2-dimethylamino-1-phenyl-1-benzyl-1-propanol, and 1-(N,N-dimethylamino)-1-phenyl-propyramine, or -NR⁴Y.

The carboxylic acids forming the organic nonlinear optical substances of this embodiment can be represented by the formula (VI):

$$(z^5 \rightarrow CH = CH \rightarrow CH = C - COOH)$$
 (VI)

Examples of such carpoxylic acid may include for example, 3-(5-membered ring derivatives with heterolaticms) substituted 2-cyanopropenoic acids such as

3-(3-thienyl)-2-cyanopropenoic acid. 3-(2-pyrrolyl)-2-cyanopropenoic acid. 3-(2-pyrrolyl)-2-cyanopropenoic acid. 3-(3-pyrrolyl)-2-cyanopropenoic acid. 3-(3-furyl)-2-cyanopropenoic acid. 3-(3-indolyl)-2-cyanopropenoic acid. 3-(N-methyl-2-cyanopropenoic acid. 3-(N-methyl-2-cyanopropenoic acid. 3-(N-methyl-2-cyanopropenoic acid. 3-(N-methyl-2-cyanopropenoic acid. 3-(N-methyl-2-cyanopropenoic acid. 3-(N-n-butyl-2-cyanopropenoic acid. 3-(N-n-butyl-2-cyanopropenoic acid. 3-(N-n-butyl-2-cyanopropenoic acid. 3-(S-nitro-2-furyl)-2-cyanopropenoic acid. 3-(5-nitro-2-thienyl)-2-cyanopropenoic acid.

5-(5-membered ring derivatives with hetero atoms) substituted 2-cyano-2.4-pentadierioic acids such as 2-cyano-5-(3-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(2-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(2-pyrrolyl)-2,4-pentadienoic acid, 2-cyano-5-(3-furyl)-2,4-pentadienoic acid, 2-cyano-5-(3-indolyl)-2,4-pentadienoic acid, 2-cyano-5-(3-indolyl)-2,4-pentadienoic acid, 2-cyano-5-(N-methyl-2-pyrrolyl)-2,4-pentadienoic acid, 2-cyano-5-(N-methyl-3-pyrrolyl)-2,4-pentadienoic acid, 2-cyano-5-(N-ethyl-2-pyrrolyl)-2,4-pentadienoic acid, 2-cyano-5-(N-n-butyl-2-pyrrolyl)-2,4-pentadienoic acid, 2-cyano-5-(N-n-butyl-2-pyrrolyl)-2,4-pentadienoic acid, 2-cyano-5-(N-n-butyl-2-pyrrolyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-2-furyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-3-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-2-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-3-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-2-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-3-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-2-thienyl)-2,4-pentadienoic acid, 2-cyano-5-(5-nitro-2

5-(5-membered ring derivatives with hetero atoms) substituted 2-cyano-2,4,6-heptatrienoic acids such as 2-cyano-7-(3-thienyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(2-thienyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(2-thienyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(2-furyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(2-indolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(2-indolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(3-furyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(2-indolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(3-indolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(N-methyl-3-pyrrolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(N-methyl-3-pyrrolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(N-n-butyl-2-pyrrolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(N-n-butyl-2-pyrrolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(N-n-butyl-2-pyrrolyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(S-nitro-2-furyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(5-nitro-2-furyl)-2,4,6-heptatrienoic acid, 2-cyano-7-(5-nitro-2-furyl)-2,4,6

EXAMPLE

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The present invention will now be further illustrated by, but is by no means limited to, the following Examples.

Synthetic Example 1

Synthesis of 2-cyano-5-(4-dimethylaminophenyl)-2,4-pentadienoic acid (1)

To 100 mt of an aqueous solution of 2.55 g of sodium hydroxide 5.97 g of methyl cyanoacetate was added and further 9.55 g of p-dimethylaminocinnamovt aldehyde was added under stirring, followed by heating to 85°C with continuous stirring for 40 hours. After completion of the reaction, 50 mt of 12 N hydrochloric acid was added, and a solid was recovered. The solid was subjected to recrystallization from methanol, repeated twice, to give 6.38 g of the desired product, m.p. 218 - 219°C. The elemental analysis of C. 68.40°c, N. 11.30°c coincided well with the calculated values of C. 69.36°c, H. 5.84°c, N. 11.56°c. IR absorption spectrum; presence of CN group at 2216 cm⁻¹, COOH group at 1673 cm⁻¹, benzene ring and conjugate double bond at 1615, 1586, and 1551 cm⁻¹ being recognized. In NMR spectrum, absorption by methyl group was recognized at 3.08 ppm, and AB type absorption based on benzene ring at 6.80 and 7.60 ppm, λmax was found to be 440 nm.

Synthesis Example 2

Synthesis of 2-cyanci-3-(4-dimethylaminophenyl)-2-propenoic acid (2)

In 400 ml of an aqueous solution of 13.77 g of sodium hydroxide, 34.80 g of methyl cyanoacetate was dissolved and 34.01 g of p-dimethylamino benzaldehyde was then added under a nitrogen atmosphere, followed by adding 200 ml of ethanol to obtain a uniform solution. Under reflux, the stirring was continued for 51 hours and the reaction mi-ture was added to 12 N hydrochloric acid to obtain the precipitates. The resultant solid was repeated twice to recrystallize from a methanol ethanol mixture to obtain 13.51 g of the needles

NMR spectrum: methyl group at 3.08 ppm, doublet benzene ring at 6.84 - 6.82 ppm and 7.93 - 7.95 ppm, and -CH = group at 8.25 ppm

Yield: 37%

m.p.: 226 - 228 ° C

Elemental analysis: C 66.82%. H 5.56%, N 12.76% (Calc. C 66.14%, H 5.60%, N 12.96%) λmax in ethanol: 399 nm

Synthesis Example 3

20 Synthesis of 2-cyano-5-(4-methoxyphenyl)-2,4-pentadienoic acid (3)

The desired compound was prepared in the same manner as in Synthesis Example 1 by using 16.2 g of p-methoxycinnamic aldehyde having a melting point of 45.5 °C, which was obtained from p-methoxystyrene and phosphorus trichloride according to a method disclosed in J. Amer. Chem. Soc., 78, 3209 (1956). 4.8 g of sodium hydroxide and 11.3 g of methyl cyanoacetate. The product was recrystallized from ethanol to obtain the needles having a melting point of 240 °C at a yield of 69%

Elemental analysis: C 68.11%, H 4.81%, N 6.10% (Calc. C 68 10%, H 4.85%, N 6.11%)

NMR spectrum: methyl group at 3.83 ppm, doublet benzene ring around 7.02 - 7.64 ppm, and -CH = group at 7.09, 7.59 and 8.06 ppm.

λmax in ethanol: 372 nm

Synthesis Example 4

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Synthesis of 2-cyano-3-(3,4-methylenedioxyphenyl)-2-propenoic acid (4)

To an aqueous solution containing 14.20 g of sodium hydroxide and 33.75 g of methyl cyanoacetate, 30.32 g of 3,4-(methylenedioxy)benzaldehyde was added, followed by stirring at 95°C for 16 hours. After completion of the reaction, an aqueous diluted hydrochloric acid solution was added to obtain a pale yellow solid. The resultant solid was recrystallized from ethanol to obtain a crystal having a melting point of 233°C.

Elemental analysis: C 61.01%, H 3.21%, N 6.37% (Calc. C 60 83%, H 3.26%, N 6.45%)

Infrared spectrum: Absorption by -CN at a wavelength of 2224 cm⁻¹, -COO- at 1677 cm⁻¹, conjugated system at 1575 cm⁻¹ and 1293 cm⁻¹

NMR spectrum: -CH, - at 6.19 ppm(s), -CH= at 8.22 ppm(s), and -H based upon a benzene ring at 7.12, 7.63, and 7.68 ppm

Synthesis Example 5

Synthesis of 2-cyano-3-(3.4-dimethoxyphenyl)-2-propensic acid (5)

To 150 ml of an aqueous solution of 9.19 g of sodium hydroxide, 20.50 g of methyl cyanoacetate was added and, under stirring, 25.38 g of 3,4-dimethoxy-benzaldehyde was further added, followed by heating at 85°C for 40 hours under stirring. After completion of the reaction, the reaction mixture was added to 50 ml of 12 N hydrochloric acid to recover the resultant solid. The solid was repeated twice to recrystallize from ethanol to obtain 19.84 g of the desired compound.

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M.p. 206.13 °C

1573, and 1512 \mbox{cm}^{-1} and presence of conjugated double bond.

NMR spectrum. Absorption by methyl group at 3.97 - 4.01 ppm, ABX type absorption based upon benzene ring at 7.00, 7.55, and 7.88 ppm.

xmax in ethanol, 353 nm

Synthesis Example 6

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Synthesis of 2-cyano-3-(2.4-dinitrophenyl)-2-propenoic acid (6)

The desired compound (6) was obtained in the same manner as in Synthesis Example 5, except that 2,4-dinitrobenzaldehyde was used instead of 3,4-dimethoxybenzaldehyde.

M.p.: 210 ° C

Elemental analysis: C 46.00%. H 1.98%, N 16.03% (Calc. C 46.53%, H 1.92%, N 15.97%

15 Synthesis Example 7

Synthesis of 2-cyano-5-(3,4-dimethoxyphenyl)-2.4-pentadienoic acid (7)

The desired compound (7) was obtained in the same manner as in Synthesis Example 1, except that 2-20 (3,4-dimethoxyphenyl)-1-formyl 1-propenoic acid obtained from the reaction of 3,4-dimethoxy benzaldehyde and phosphorus trichloride.

M.p.: 190 ° C

Elemental analysis: C 64.70%, H 5.15%, N 5.62% (Calc. C 64.85%, H 5.06%, N 5.40%)

25 Synthesis Examples 8 - 14

The compounds (8) - (14) shown in Table 1 were synthesized from the corresponding aldehydes and methyl cyanoacetate in the manner described above.

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Table 1: Synthesis of Aromatic Derivative

Com- pound	Structure	m.	.p. (°C)			λma x *	
No.	Elemental	analysis	(C, H,	N)(F	: Foun	.d/C: Ca	lcd)
8	p-NO ₂ -C ₆ H ₄ -CH=C(CN)CO F 5	ОН	208			302nm	
	2 6 4 F 5	4.91 2.9	33 12.80	/ C	55.05	2.75	12.3
9	D+CHO-C_HCH=C(CN)	соон	229			320nm	
	P-CH ₃ -O-C ₆ H ₄ -CH-C(CN)	5.10 4.6	6.71	/ C	65.02	4.96	6.8
10	H-C.HCH=C(CN)COOH		210			295nm	
	H-C ₆ H ₄ -CH=C(CN)COOH F 6	9.58 4.3	8.03	/ C	69.35	4.08	8.0
11	C_H_=CH=CH=CH=C(CN)	соон	212			320nm	
	C ₆ H ₅ -CH=CH-CH=C(CN) F 7	2.50 4.6	50 7.01	/ C	72.34	4.56	7.0
12	C_HCH=CH-CH=CH-CH	-C(CN)C	он 238			360nm	
	C ₆ H ₅ -CH=CH-CH=CH-CH F 7	4.75 4.8	38 6.35	/ C	74.64	4.93	6.2
13	m-CH ~O-C .HCH=C(CN)	соон	166			296nm	
	m-CH ₃ -O-C ₆ H ₄ -CH=C(CN) F 7	0.69 6.6	57 12.37	/ C	71.18	6.88	12.4
14	p-C . H . O-C .HCH=C(C	N)COOH	82			-	
_ :	p-C ₁₀ H ₂₁ O-C ₅ H ₄ -CH - C(C F 7	3.55 8.5	55 4.15	/ C	73.43	8.53	4.0

*1: determined in methanol

35 Synthesis Example 15

Synthesis of trans,trans,trans,2-cyano-7-(n-pentyl)-2.4,6-heptatriene-1-carboxylic acid (15)

To 150 ml of an aqueous solution containing 6.87 g of sodium hydroxide and 16.40 g of methyl cyanoacetate 14.85 g of trans,trans-2.4-decadenal was added, followed by heating at 100°C for 16 hours under stirring. After completion of the reaction, an excess amount of an aqueous hydrochloric acid solution was added thereto to obtain a viscous solid. The resultant solid was recrystallized from n-hexane to obtain the crystal having a melting point of 98 - 102°C.

Elemental analysis: C 70.00%, H 7.75%, N 6.27% (Calc. C 71.19%, H 7.83%, N 6.39%)

IR spectrum -CN at 2211 cm⁻¹, -COO- at 1609 cm⁻¹, conjugated system at 1561 cm⁻¹ and 996 cm⁻¹ NMR spectrum: -CH = CH- at 6.25 - 7.95 ppm, long CH; - and CH; - at 0.85 - 2.2 ppm, integrated intensity was well coincided

Synthesis Example 16

Synthesis of trans.trans.2-cyano-5-(n-heptyl)-2.4-pentadienoic acid (16)

The desired compound was synthesized and purified in the same manner as in Synthesis Example 15, except that trans 2-decenal was used as a starting material. Thus crystals were obtained.

Elementa analysis C 71 20%. H 8.90% N 6 17% (Calc C 70.55% H 8.67% N 6 33%)

Synthesis Example 17

Synthesis of 2-cyano-3-(2-thienyli-2-propenoic acid (17)

To 160 ml of an aqueous solution containing 20.97 g of sodium hydroxide and 46.11 g of methyl cyanoacetate, 40.08 g of thiophene 2-carboxyaldehyde was added, followed by heating at 90°C for 9 hours under stirring. After completion of the reaction, an excess amount of hydrochloric acid was added and the resultant solid was recovered. The solid was recrystallized from ethanol to obtain the needles.

M.p.: 234 * C

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Elemental analysis: C 53.63%, H 2.69%, N 7.80%, S 17.70% (Calc. C 53.61%, H 2.82%, N 7.82%, S 17.89%)

NMR spectrum: proton of thiophen ring at 7.34, 8.02 and 8.17 ppm, proton at p-position at 8.55 ppm λmax in ethanol: 335 nm

15 Synthesis Example 18

Synthesis of 2-cyano-3-(3-thienyl)-2-propenoic acid (18)

The desired compound (18) was prepared in the same manner as in Synthesis Example 17, except that thiophene 3-carboxyaldehyde was used instead of thiophene 2-carboxyaldehyde.

M.p.: 211 °C

Elemental analysis: C 53.73%, H 2.71, N 7.73%, S 17.52% (Calc. C 53.61%, H 2.82%, N 7.82%, S 17.89%)

25 Synthesis Example 19

Synthesis of 2-cyano-3-(2-pyrrolyl)-2-propenoic acid (19)

After 36.93 g of methyl cyanoacetate and 16.94 g of sodium hydroxide were dissolved in 260 ml of water, 23.80 g of pyrrole 2-carboxyaldehyde was added thereto, followed by heating at 95 °C for 30 hours under stirring. Thereafter, hydrochloric acid was added and the resultant solid was recrystallized from a mixed solvent of ethanol/methanol to obtain the crystal having a melting point of 213 °C.

Elemental analysis: C 59.34%, H 3.82%, N 17.26% (Calc.: C 59.25%, H 3.73%, N 17.28%)

35 Synthesis Example 20

Synthesis of 2-cyano-3-(2-furyl)-2-propenoic acid (20)

The desired compound (20) was obtained in the same manner as in Synthesis Example 17, except that 40 furfural was used.

M.p.: 219 ° C

Elemental analysis: C 59.02%, H 2.95, N 8.53% (Calc.: C 58.89%, H 3.10%, N 8.59%) λmax in ethanol: 330 nm

45. Synthesis Example 21

Synthesis of 2-cyane-5-(2-furyl)-2,4-pentadienoic acid (21)

The desired compound (21) was prepared in the same manner as in Synthesis Example 17 except that 50 24.7 g of 3-(2-furyl) acrolein was used.

The melting point of the resultant compound was 220°C and the structure thereof was confirmed by the elemental analysis value and NMR spectrum. The \(\text{\lambda} \text{max} \) in ethanol was 368 nm.

Synthesis Example 22

Synthesis of 2-cyano-7-(2-turyl)-2 4 6-heptatriene-1-carboxylic acid (22)

The desired compound (22) was prepared in the same manner as in Synthesis Example 21 except that the aldehyde obtained from the oxidation reaction with phosphorus oxytrichioride mentioned in Synthesis Example 3. The transistructure of the resultant was confirmed by an NMR spectrum.

Synthesis Example 23

Synthesis of 2-cyano-3-(3-indoly))-2-propenoic acid (23)

The desired compound (23) in the form of a pale yellow flake crystal was obtained at a yield of 33.5% in the same manner as in Synthesis Example 1 by using 21.34 g of indole 3-carboxyaldehyde, 9.47 g of sodium hydroxide and 23.46 g of methyl cyanoacetate.

M.p. 230 °C

Elemental analysis: C 68.33%, H 3.77%, N 13.29% (Calc., C 67.92%, H 3.80%, N 13.20%) \(\lambda\) max in ethanol, 378 nm

20 Evaluation method of intensity of second harmonic generation

According to the method as described in S. K. Kurtz et al., Journal of Applied Physics (J. Appl. Phys.). Vol. 39, p. 3798 (published in 1968), the generation of the second harmonic wave was measured for the powder of the compound of the present invention. As the incident ray source, a beam of 1.06 μ of Nd:YAG laser (2 W/2 KHz pulse) was employed and irradiated on the powdery sample filled in a glass cell. The incident light was filtered by a filter and the green light generated at an incident angle of 55° from the normal direction was detected to obviate the effect from the strength of the incident light. As the sample for Control, urea powder or m-nitroaniline powder having a particle size of 50 to 90 μ m, which was previously powdered, followed by sieving, was employed. Regarding the resistance to a laser beam, the laser beam was irradiated to the sample and the deformation in the appearance was visually observed. Generally, the determination of the principal characteristics was carried out at a non-focus point because the intensity of the laser beam was strong.

Example 1

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A 2.39 g amount of the carboxylic acid compound (1) obtained in the Synthesis Example 1 was dissolved in 150 ml of tetrahydrofuran and, to the resultant solution, 1.18 g of L-(-)-1-phenylethylamine was added under stirring. Momentarily, the precipitate was generated, which was filtered to recover 3.16 g of a reddish orange solid. The solid was recrystallized from ethanol/methanol mixture to obtain 2.16 g of needles. The elemental analysis value of this product was as follows.

C: 72 70% (Calc. *72.09%), H: 6.68% (Calc. *6.95%), N: 11.63% (Calc. *11 56%)

IR spectrum: carboxylate at 2400 - 3200 cm-

In the case of the carboxylic acid (1), the absorbance of the COOH group at 1673 cm⁻¹ was shifted to about 1620 cm⁻¹ to exhibit the generation of the salt formation.

The NMR spectrum exhibited an absorbance by a methyl group at 2.95 ppm, benzene at 6.74 - 7.50 ppm, and methyl group of 1-phenylethylamine at 1.50 ppm. The relative ratio of the methyl group absorbance intensity was 2.1, indicating the formation of a salt of carboxylic acid amine = 1.1. The equivalent molar ratio was also confirmed from the elemental analysis. The λ max of this salt in ethano was 420 nm, which was changed to the low wavelength side by 20 nm when compared with that of the corresponding carboxylic acid (1). The melting point was 188°C and the degree of the optical rotation {a}o at Na-D ray in methanol was -5° (c = 0.597). When the powder was irradiated by a 1.06 μ light of Nd-YAG laser, the intensity of the second harmonic generation was about 3 times of that of m-nitroaniline.

Example 2

The carboxylic acid (12) prepared in the Synthesis Example 12 was used to form a salt with L-(-)-1-phenylethylamine in a THF solution in the same manner as in Example 1. The crystals were precipitated with the elapse of time. The resultant crystal was recrystallized from a mixed solvent of methanoliethanol to obtain a pale yellow crystal having a melting point of 172°C.

The elemental analysis of the resultant crystal is C: 75.98%, H 6.18%, and N 8.06%, which was well coincided with the calculated value of C: 76.26%, H: 6.41%, and N: 8.09% in terms of a 1:1 salt of the carboxylic acid (12) and phenylethylamine

IR spectrum showed broad carboxylate absorption at 2400 - 3200 cm⁻¹, and the absorption of COOH group at 1673 cm⁻¹ in the carboxylic acid (12) was shifted to about 1620 cm⁻¹ to reveal the formation of the salt.

The NMR spectrum gave an integrated intensity suggesting the formation of carboxylic acid amine of 1.1. The degree of the optical rotation $[\alpha]_D$ at Na-D ray in methanol was +0.97 degree (c = 0.597). The λ max in ethanol of the resultant salt was 355 nm, which was changed to the low wavelength side by 5 nm when compared with that of the corresponding carboxylic acid. When the powder was irradiated by a 1.06 μ light of Nd-YAG laser, the intensity of the second harmonic generation was about 1.8 times of that of m-nitroaniline.

20 Examples 3 - 12

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Optically active amine salts of various carboxylic acid compounds were obtained in the same manner as in Example 1, and their second harmonic generating abilities were determined. The results are shown in Table 2.

Table 2

Example No.	Carboxylic acid amine	m.p. (*C)	Elemental analysis (Found/Calc.)	Degree of optical rotation [a] _D	λmax	SHG * generating ability
3	11 PEA	147	C:74.92%, H:6.14%, N: 8.77% (C:74.96%, H:6.30%, N: 8.74%)	-2.01	325 nm	1.2
4	2 PE A	177	C:70.69%, H:6.67%, N:12.37% (C:71.18%, H:6.88%, N:12.46%)	-0.33	385 nm	0.4
5	9 PEA	136	C:69.96%, H:6.04%, N: 8.61% (C:70.34%, H:6.23%, N: 8.64%)	-0.88	320 nm	0.3
6	10 PEA	164	C:72.82°c, H:5.98%, N: 9.58°c (C:74.45°c, H:6.26%, N: 9.65°c)	-0.73	288 nm	0.1
7	13 PEA	149	C:70.41°E, H:6.23°6, N: 8.62% (C:70.34°6, H:6.23°6, N: 8.62°6)		285 nm	0.3
8	5 PEA	207	C:67.74%, H:6.12%, N: 7.94% (C:67.77%, H:6.27%, N: 7.91%)		333 nm	0.2
9	4 PEA	176	C 67 79°c, H:5.37°c, N: 8.21°c (C:67.43°c, H:5.37°c, N: 8.28°c)		337 nm	03
10	8 PEA	154	C 63 31%, H:4.78?%, N 12.25% (C 63.70%, H:7.94%, N 12.38%)	+ 2 35	302 nm	0.1
11	23 PEA	175	C: 71.90°c, H.5.64%, N:12.52% (C: 72.04°c, H:5.76°c, N:12.61%)		358 nm	0.1
12	20 PEA	150	C 67.90°c, H:5.64%, N: 9.52% (C: 67.58%, H:5.68%, N: 9.85%)		320 nm	0.2

^{* (}relative to m-nitroaniline (m-NA))

r, r

Example 13

A 0.67 g amount of the carboxylic acid 4 obtained in Synthesis Example 4 was dissolved in 7 ml of tetrahydrofuran, followed by adding 0.92 g of optically active R-(-)-1- $(\alpha$ -naphthyliethylamine. The resultant solid was filtered and recrystallized from ethanol. The melting point was 1.71 °C and the ratio of the integrated intensity of the absorption peaks of the carboxylic acid (4) component to (α -naphthyl)ethylamine component such as methyl group obtained from the NMR spectrum was 1:1.

The elemental analysis was C 71.55%. H 5.20%. N 7.21%, which was coincided with the calculated value, C 71.55%. H 5.26%, N 7.20% which was based upon the formation of a 1:1 salt of the carboxylic acid and the amine.

When the resultant crystal was finely powdered and the generation of the second harmonic generation was determined, the intensity of the emission is about 1.5 times of that of the urea.

Example 14

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A 0.93 g amount of the carboxylic acid (1) obtained in Synthesis Example 1 was dissolved in 10 ml of tetrahydrofuran, followed by adding 0.70 g of S-(-)-1-(α -naphthyl)ethylamine thereto. The resultant solid was recovered and washed thoroughly with tetrahydrofuran. After drying, the crystal having a melting point of 171 °C. The degree [α]_D of the optical rotation of the sample in methanol was -30.0 degree (c = 0.04) and λ max was 421 nm. The maximum wave length was approximately coincide with that of Example 1. The second harmonic generation generating ability of the crystal was about 3.9 times of that of urea. Furthermore, when the crystalline powder was exposed to the laser beam for a long time, apparent damage was not observed.

25 Examples 15 - 19

Various optically active amine salts were prepared in the same manner as in Example 14 and the second harmonic generating abilities were determined.

The results are shown in Table 3.

Table 3

	Example	Carboxylic acid	Optically active amine	SHG generating ability *1
	15	Synthesis Example 5	1-Phenyl-2-methylethylamine	1.5
	16	Synthesis Erample 7	1-Phenylethylamine	5.0
- 1	17	Synthesis Example 7	1-(a-naphthyl)ethylamine	3.5
	18	Synthesis Example 5	1-Phenylethylamine	2.0
	19	Synthesis Example 5	1-Phenyl-2-aminopropane	2.5

11: Based on urea powder

Example 20

A 1.54 g amount of the thiophene-containing carboxylic acid (17) obtained in Synthesis Example 17 was dissolved in 40 ml of tetrahydri furan, followed by adding 1.46 g of optically active R-G-1-phenylethylamine. The precipitated solid was recovered and recrystallized from ethanol to obtain the crystal having a melting point of 171 °C (decomposition).

The elemental analysis values were C 63.85%, H 5.15%, N 9.30%, S 10.40%, which were well coincide with the calculated values, C 63.97%. H 5.38%. N 9.33%, S 10.17% based upon a 1.1 amine salt of the carboxylic acid. The integrated intensity ratio of the absorption peak of the carboxylic acid of Synthesis Example 17 and phenethylamine from the NMR spectrum. The maximum absorption wavelength was 322 nm. When the second harmonic generation was determined after the crystal was finely powdered, the green emission having the ability of about twice of that of urea was observed

E>ample 21

A 1.51 g amount of the 3-substituted thiophene carboxylic acid (18) obtained in Synthesis Example 18 was dissolved in 20 ml of tetrahydrofuran, followed by adding 1.20 g of R-i-)-1-phenylethylamine. The precipitated solid was recovered and recrystallized from ethano, after drying, to obtain the crystal having a melting point of 171 °C (decomposition)

The elemental analysis values were C 63.82%, H 5.07%, N 9.31%, S 10.57%, which were well coincide with the calculated values, C 63.97%, H 5.38%, N 9.33%, S 10.17% based upon a 1:1 amine salt of the carboxylic acid. The integrated intensity ratio of the absorption peak of the carboxylic acid of Synthesis Example 18 and phenethylamine from the NMR spectrum. The maximum absorption wavelength was 322 nm. The second harmonic generating ability of the crystal was 3 times of that of urea and the maximum atsorption wavelength of the sample in ethanol was 322 nm.

Example 22

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A 0.99 g amount of the carbo-ylic acid (21) obtained in Synthesis Example 21 was dissolved in 40 ml of tetrahydrofuran, followed by adding 0.78 g of optical active R-(-)-1-phenylethylamine thereto

After n-hexane was then added, the precipitated solid was recovered. The solid was recrystallized from ethanol to obtain the crystal having a melting point of 121 °C (decomposition). The elemental analysis data were C 69.55%, H 5.95%, and N 9 00%, which was well coincided with the calculated values of the optically active amine salt of the synthesized carboxylic acid (21). The emission having an intensity of about 5 times of that of urea was observed by an Nd-YAG laser beam. The maximum absorption wavelength of the sample in ethanol was 350 nm.

25 Example 23

A 3.20 g amount of the carbo-ylic acid (12) obtained in Synthesis Example 12 was dissolved in 50 ml of tetrahydrofuran, followed by adding 2.50 g of R-(-)-2-amino-1-butanol thereto.

The precipitation was momentarily generated and, after filtering, 3.00 g of yellow solid was recovered a mixed solvent of ethanol/methanol to obtain 2.1 g of the needle having a melting point of 187 °C. The optical rotation degree in methanol with an Na-D ray was 16 degree. The maximum absorption of the sample in methanol was 355 nm amine salt of the synthesized carboxylic. The second harmonic generating ability of the powder was 33 times of that of urea.

35 Examples 24 - 42

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The salts of various carboxylic acids and optically active alcohol amines were carried out in the same manner as in Example 23 and the second harmonic generating abilities of the resultant crystals were determined.

The results are shown in Table 4.

Table 4

£	Example	Carboxylic acid	Optically active amine base	SHG generating ability *1
÷ ;	24	23	1-Amino-2-propano!	
	25	1	2-Amino-1-butanol	6.9
	26	2	11	1.8
	27	10	"	1.2
70	28	2	1-Amino-2-propanol	30.0
	29	10	ti .	1.1
	30	1	п	26.0
	31	12	n	1.1
	32	2	2-Amino-1-propanol	0.5
15	33	10	n e	0.2
	34	12	2-Amino-1-(p-nitrophenyl)-1,3-propane diol	0.7
	35	12	2-Dimethylamino-1-phenyl-1-phenyl-1-benzyl-1-propanol	0.6
	36	17	2-Amino-1-butanol	3.0
	37	17	2-Amino-1-propanol	1.5
20	38	4	"	1.3
	39	20	"	3.8
	40	20	2-Amino-1-butanol	5.0
	41	21	"	3.6
	42	19	н	3.2

'1: Based upon urea powder

Example 43

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A 0.89 g amount of dimethoxy substituted conjugated carboxylic acid (7) obtained in Synthesis Example 7 was dissolved in 10 ml of THF, followed by adding 0.42 g of dextrorotatory R-(-)-2-amino-1-butanol. The resultant precipitates was filtered and recrystallized from ethanol to obtain the white crystal having a melting point of 130.5 °C. The NMR spectrum of the resultant solid revealed the integrated intensity, which shows the formation of 1:1 by mole salt of the corresponding carboxylic acid and amine. The crystal was finely powdered and the second harmonic generating ability was determined. As a result, the emission capability thereof was 5.8 times of that of urea. The maximum absorption of the sample in ethanol was 370 nm. The powder of 2-methyl-4-nitroaniline used as a control was melted and carbonized under the determination conditions of the second harmonic generating ability, whereas the change in the emission capability of the present sample with the elapse of time was not observed and the damage resistance of the present sample against light was good.

Example 44

The amine salt was formed in the same manner as in Example 43, except that R-(-)-1-amine-2-propanol was used as the optically active amine. The second harmonic generating ability of the resultant salt was about 4 times of that of uroa and the ability was not changed with the clapse of time. Thus, it has been observed that a high resistance to the light damage was good.

50 Example 45

The salt was formed in the same manner as in Example 43, except that the dimethoxy compound (5) obtained in Synthetic Example 5 as the carboxylic acid. The emission capability of the resultant salt was not changed when the sample was exposed to a laser beam for a long time.

Example 46

A 2.30 g amount of L-phenylalanine ethyl ester hydrochloride was suspended in 50 ml of other, followed by adding thereto 0.96 g of triethyl amine and further 30 ml of water, ten ml of the supernatant other phase was recovered and this solution was added to a previously prepared solution of 0.26 g of the above-mentioned carboxylic acid (1) in 6 ml of THF. The needles were obtained with the elapse of time. The decomposition of this crystal was 180 °C. When the second harmonic generating ability of this crystal was determined, the intensity was about 15 times of that of urea.

to Example 47

A 3.30 g amount of L-valine methyl ester hydrochloride was suspended in 50 ml of ether, followed by adding thereto 1.89 g of triethyl amine to obtain an ether solution of the L-valine methyl ester. This solution was added to a solution of 0.12 g of the above-mentioned carboxylic acid (2) in 10 ml of THF to obtain the needles. When the second harmonic generating ability of the crystal powder was determined, the intensity was 3 times of that of urea.

E> amples 48 - 64

20 The amine salts of α-amino acid esters of various carboxylic acids were obtained in the same manner as in Examples 1 and 2 and the second harmonic generating abilities were determined.
The results are shown in Table 5.

Table 5

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-	v

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Example	Carboxylic acid	Optically active amine base	SHG generating ability *1
48	10	L-Phenylalanine ethyl ester	16
49	12	#	10
50	8	н	5
51	10	"	7
52	13	*	5
53	15	*	4
54	19	*	4
55	1	L-Valine methyl ester	12
56	11	-	15
57	14	-	4
58	12	**	18
59	16	#	3
60	1	D-Phenylglycine methyl ester	10
61	9	,	5
62	7		11
63	1	α-N-Benzyol-L-aginine ethy! ester	8
64	7	,	3

*1 Based on urea

50 Example 65

Synthesis of S-(-)-phenethyl 2-cyano-5-phenyl-2.4-pentadienocarboxamide (24)

A 3.3 g amount of an acid chloride of the compound (11) obtained by thermally treating the compound (11) with thionyl chloride was added, under vigorous stirring, to 20 ml of dioxane containing 1.8 g of S-(-i-phenethyl amine and 1.5 g of triethyl amine. After stirring at room temperature for further 3 hours, the

Elemental analysis C 79.50% H 6.05% N 9.30% (Calc C 79.44% H 6.00% N 9.26%)

IR absorption spectrum -NH- group at 3364 cm⁻¹. CN group at 2216 cm⁻¹ amine I II at 1649 cm⁻¹ and 1522 cm⁻¹

Amax in dioxane 336 nm

When the second harmonic generating ability of the crystal was determined, the intensity was about 11 times of urea and there are no substantial deformation, when the sample was irradiated for a long time.

Example 66

Synthesis of S-(-)-phenethyl 2-cyano-7-phenyl-2,4.6-heptatriene-1-carboxamide (25)

A 1.79 g amount of the acid chloride of the compound (11) (m.p. 143 °C) obtained by thermally treating the compound (11) with thionyl chloride was added, under vigorous stirring, to 30 ml of THF containing 1.06 g of S-(-)-phenethyl amine and 0.70 g of pyridine dissolved therein. After completion of the reaction, a large amount of water was added to the reaction product and the resultant precipitates were filtered, followed by recrystallizing, to obtain 1.6 g of a crystal having a melting point of 128 °C.

Elemental analysis: C 80.74%, H 6.20%, N 8.66% (Calc. C 80.44%, H 6.15%, N 8.53%)

IR absorption spectrum. -NH- group at 3360 cm⁻¹, -CN group at 2216 cm⁻¹, presence of amine I, II at 1649 cm⁻¹ and 1522 cm⁻¹.

λmax in ethanol: 370 nm

NMR spectrum: the methyl group of phenethyl group at 1.57 ppm, -CH = at 7.99 ppm, 6.79 - 7.05 ppm, 6.82 ppm (Thus, the structure of the desired compound was confirmed)

When the second harmonic generating ability of the crystal was determined, it was observed that the intensity was 1.1 times of urea and there are no substantial deformation after a long time irradiation of a laser beam.

Example 67

Synthesis of S-(-)1-(α -naphthyl)ethyl 2-cyano-7-phenyl-2.4.6-heptatriene-1-carboxamide (26)

To a dry THF solution of an equivalent mixture of the compound (12) and S-(-)1-(a-naphthyl)ethylamine, dicyclohexylcarbo diimide was added, followed by stirring for one night. The precipitated dicyclohexyl urea was recovered by filtration and the mother liquor was concentrated, followed by recrystallizing from a mixture of ethanol/methanol to obtain a white solid having a melting point of 100 °C.

IR absorption spectrum: -NH- group at 3360 cm $^{-1}$, CN group of 2220 cm $^{-1}$, presence of amide I, II at 1650 cm $^{-1}$ and 1522 cm $^{-1}$

When the second harmonic generating ability of the crystal was determined, the intensity was 4 times of that of urea and there are no substantial deformation after a long time irradiation of a laser beam.

40 Examples 68 - 72

Various optically active amides shown in Table 6 were obtained from various carbokylic acid and optically active acid amide

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		Am1de
	٠	T V
		e VC
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	:	Optically
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		Table

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Example Carboxylic acid comp 68		Amine	Canthonic		
3 P 2 C ₆ 3 BA:	Carboxylic acid component	component	synthesis method	(°C)	SHG
PEA:	p-N0 ₂ -C ₆ H ₄ -CH-C(CN)COOH	PRO	DCC	135	3
PEA:	р-сн3-0-св4-сн-с(си)соон	S-(-)PEA	CL	126	S
C C C C C C C C C C C C C C C C C C C	C(CN)COOH	R-(-)-sBA	CL	98	3
PEA:	р-СН ₃ -0-С ₆ Н ₄ -СН-СН-СН-С (СN)СОЭН	•		140	S
) PEA:	C ₆ H ₅ -CH-CH-CH-CH-C(CN)COOH	S-(-)PEA	DCC	158	S
) PEA:	101				
) s BA:	S-(-)-phenethylamine				
	R-(-)-2-amino-1-butanol				
	Dicyclohexyl carbodiimide method (Example 67)	od (Example	(2)		
	Acid chloride method (Example 66)	(99			
	ssion				
S: Strong emission	mission				

Claims

1. An organic nonlinear optical substance having the formula (I):

$$A \leftarrow CR^{1}=CH \rightarrow CH = C - C - B \qquad (I)$$

$$CN = C$$

wherein

R' represents -H or -CH₃; n is 0, 1, or 2; A represents Z'-Ar-,

 $\xi_{i,\xi}$

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$$z^2$$
 z^3
Ar-, z^2 -, or (z^5)
 z^4

wherein Ar represents a 6 - 14 membered aromatic group including a heterocyclic ring or bisphenylene type ring. Z' represents H-, R' R' N-, R' 0-, R's S-, NC-, R' 0C0-, R' C00-, O_2 N-, R' R' 2NOC-, R' 3C0-(R''s)N-, or R''s-; Z'', Z'', and Z''s independently represent H-, a C--C_k alkyi-, R''s 0-, R'''R''s N, R''s S-, O_2 N-, or two R''s being, in combination, R's CH-, : R' represents H- or a C₁-C₁₂ alkyl; R's to R''s independently represent H-, or a C--C_k hydrocarbon residue, Z's independently represents H-, a C--C_k saturated hydrocarbon residue, O_2 N-, R'''s N-, wherein R''s to R''s independently represent H or a C--C_k saturated hydrocarbon residue, X represents -S-, -0-, or O_2 NR''s r is 0 cr an integer of 1 to 3; and R''s represents H or a hydrocarbon group having 1 to 8 carbon atoms;

B represents -OH+Amine* where Amine represents an optically active amine; -NR4Y where R4 represents -H or a single bond, Y represents -(CH₂)_p CQ¹Q²Q³ where p is 0 or 1; Q¹, Q², and Q³ are different and represent -H, a C·-C₅ alkyl, phenyl, naphthyl, -OH, -CH₂OH, -COOR²5, -CNR²6 R²7, a residue of an α -amino acid from which an amino group is removed, where R²5 to R²7 independently represent -H or -C₁-C₈ hydrocarbon residue; or Y represents -CQ⁴Q⁵Q⁶ where Q⁴, Q⁶ and Q₆ are different and Q⁴ and Q⁶ are as defined for Q¹, Q² and Q₃ and Q₆ represents (CH₂)_q of which one bond is linked to the bond of R⁴ where q is an integer of 1 to 4.

- 2. An organic nonlinear optical substance as claimed in claim 1, wherein in the formula (I), R¹ represents H, Z¹ represents H-, R⁵ R⁶ N-, R⁻ O-, RՑ S-, NC-, RԿ OCO-, R¹º COO-, O₂ N-, R¹¹ R¹² NOC-, R¹³ CO(R¹⁴) N-, or R¹⁵-. Ar represents an aromatic group having 5 to 14 carbon atoms, and B represents -OH• Amine* wherein Amine* represents an optically active amine selected from the group consisting of 1-phenylethylamine, 1-(α-naphthyl) ethylamine, 1-phenyl-2-methylethylamine, 1-phenyl-2-aminopropane, and brucine.
- 3. An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I), R1 represents H, Z1 represents H-, R5 R5 N-, R7 O-, R8 S-, NC-, O2 N-, R9 OCO-, R10 COO-, R112 NOC-, R13CO(R14) N-, or R15-, Ar represents an aromatic group having 6 to 14 carbon atoms, and B represents -OH- Amine* wherein Amine* represents an optically active amine selected from the group consisting of 2-amino-1-butanol, 1-amino-2-propanol, 2-amino-1-propanol, 2-amino-1-(p-nitrophenyl)-1,3-propanediol, 2-dimethylamino-1-phenyl-1-benzyl-1-propanol, and 1-(N,N-dimethylamino)-1-phenyl-propylamine.
- An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I), R¹ represents H, A represents

$$z^{2}$$
 z^{3}
 z^{4}
Au

wherein one of Z*, Z*, and Z* represents H and the remainder independently represents C+-C+calkyl, R**C+, R**R*-N-, R**S, or C+, N+, At represents an aromatic group having 6 to 14 carbon atoms, and B represents a residue of an optically active amine selected from the group consisting of 2-amino-1-butanol, 1-amino-2-propanol, 2-amino-1-propanol, 2-amino-1-phenyl-1,3-propanediol, 2-dimethylamino-1-phenyl-1-benzyl-1-propanol, and 1-(N-N-dimethylamino)-1-phenyl-propylamine.

5. An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I), R* represents H, A represents

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wherein one of Z^2 , Z^2 and Z^4 represents H- and the remainder independently represents C--C-calkyl, R*CO-, R**R*N-, R**S-, or O, N-, At represents an aromatic group having 6 to 14 carbon atoms, and B represents a residue of an optically active amine selected from the group consisting of 1-(phenyl)-ethylamine, 1-(a-naphthyl)ethylamine, 1-phenyl-2-aminopropane, and brucine.

- 6. An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I), R1 represents H. A represents Z1 Ar-, Z1 represents H-, R5R1N-, R7O-, R8S-, NC-, OzN-, R5OCO-, R10COO-, R11R12NOC-, R10CO(R14)N-, or R10-, Ar represents an aromatic ring having 6 to 14 carbon atoms, and B represents -NR4Y.
- An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I), R¹ represents H, A represents Z¹-Ar-, Z¹ represents R⁵ R⁶ N-, R² O- R⁵ S-, NC-, Rႎ OCO-, H⁻□COO-, H□COO-, H□COO-,
 - 8. An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I), R' represents H, A represents

$$z^{2}$$
 z^{3}
 z^{4}

where one of Z^2 , Z^3 , and Z^4 represents H or substituted C_1 - C_8 alkyl, the remainder of Z^2 , Z^3 , and Z^4 represents together methylene dioxy group wherein the dioxy groups are bonded to the adjacent positions of Ar, Ar represents an aromatic group having 6 to 14 carbon atoms, and B represents -OH+Amine* wherein Amine* represents an optically active amine selected from the group consisting of 1-phenylethylamine, 1- $(\alpha$ -naphthyl)ethylamine, 1-phenyl-2-methylethylamine, 1-phenyl-2-aminopropane, brucine, 2-amino-1-butanol, 1-amino-2-propanol, 2-amino-1-propanol, 2-amino-1-(p-nitrophenyl)-1,3-propanediol, 2-dimethylamino-1-phenyl-1-benzyl-1-propanol, 1-(N,N-dimethylamino)-1-phenyl-propylamine, and a residue of an optically active α -amino acid and the derivative thereof; or -NR⁴Y.

- 9. An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I), R1 represents H or CH₃, A represents R2 where R2 represents H or an alkyl group having 1 to 12 carbon atoms, and B represents -OH Amine* wherein Amine* represents an optically active amine selected from the group consisting of 1-(phenyl)ethylamine, 1-(α-naphthyl)ethylamine, 1-phenyl-2-methylethylamine, 1-phenyl-2-aminopropane, brucine; 2-amino-1-butanol, 1-amino-2-propanol, 2-amino-1-propanol, 2-amino-1-(p-nitrophenyl)-1.3-propanol, 2-dimethylamino-1-phenyl-1-benzyl-1-propanol, 1-(N.N-dimethylamino)-1-phenyl-1-benzyl-1-propanol, 1-(N.N-dimethylamino)-1-phenyl-1-benzyl-1-propanol, and a residue of an optically active α-amino acid and the derivative thereof.
- 10. An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I). R² represents
 H. A represents

$$(z^5)_r \stackrel{C}{-} \stackrel{C}{-} \stackrel{C}{-} ,$$

and B represents -OH+Amine* wherein Amine* represents an optically active amine selected from the group consisting of 1-phenylethylamine 1- $(\alpha-naphthyliethylamine 1-phenyl-2-methylethylamine 1-phenyl-2-amino-1-propanol, 2-amino-1-butanol, 1-amino-2-propanol, 2-amino-1-propanol, 2-amino-1-(p-nitrophenyli-1,3-propanediol, 2-dimethylamino-1-phenyl-1-benzyl-1-propanol, 1-(N.N-dimethylamino)-1-phenyl-propylamine, and a residue of an optically active <math>\alpha$ -amino acid and the derivatives thereof, or NR⁴Y

An organic nonlinear optical substance as claimed in claim 1, wherein, in the formula (I). R² represents
 H. A represents

$$\frac{z^2}{z^3} - Ar$$

wherein one of Z^2 , Z^3 , and Z^4 represents H- and the remainder independently represents $C \cdot - C \cdot \alpha$ alkyl, $R^{15}O^2$, $R^{17}R^{18}N^2$, $R^{19}S$, or O_2N^2 , Ar represents an aromatic group having 6 to 14 carbon atoms, and B represents -OH+Amine* wherein Amine* represents an optically active α -amino acids or the derivatives thereof.

Patentansprüche

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1. Organische nichtlineare optische Verbindung mit der Formel (I):

A
$$\leftarrow$$
 $CR^{1} = CH \rightarrow \frac{}{n}$ $CH = C - C - B$
 $CN = O$ (I)

worin R¹ Wasserstoff oder -CH₃ repräsentiert; n gleich 0, 1 oder 2 ist; A die Gruppe Z¹-Ar-,

$$z^2$$
 z^3
 z^4
Ar-, z^2 - oder (z^5)
 z^4
 z^4

repräsentiert.

worin Ar eine 6-14gliedrige aromatische Gruppe einschließlich eines heterozyklischen Rings oder Bisphenylenrings repräsentiert.

Z. Wasserstoff, R. Réin-, Rind-, Rein-, NC-, Réindon, Rindon, Rindon,

Z¹, Z² und Z⁴ unabhängig voneinander Wasserstoff, einen C--C₂-Alkylrest, R¹⁴O-- R¹¹R¹⁸N, R¹⁹S-, O₂N- oder zwei R¹⁴-Reste repräsentiert, welche kombiniert R²⁰CH ⊆ergeben;

R² Wasserstoff oder einen C·-C·2-Alkylrest repräsentiert;

 R^{∞} bis R^{20} unabhängig voneinander Wasserstoff oder einen C_1 - C_2 -Kohlenwasserstoffrest repräsentieren, Z^{ϵ} unabhängig Wasserstoff, einen gesättigten C_2 - C_2 -Kohlenwasserstoffrest, 0_0 - N_1 , R^{20} - N_2 - N_3 - N_4 - N_4 -repräsentiert, worin R^{21} bis R^{24} unabhängig voneinander Wasserstoff oder einen gesättigten C_2 - C_3 -Kohlenwasserstoffrest repräsentieren, X_1 - X_2 - X_3 - X_4 - X_4 -repräsentiert, X_4 - X_4

Wasserstoff oder eine Einfachbindung repräsentiert, Y -(CH, $_{\rm IE}$ CQ Q-Q-2 repräsentiert, wobei p gleich 0 oder 1 ist, Q1, Q2 und Q2 verschieden sind und Wasserstoff, ein C+-C+-Alkyl, Phenyl, Naphthyl, -OH, -CH_cOH_c-COOR²⁵, -CNR²⁶R-1 und einen Rest einer α-Aminosäure, von der ein Aminorest entfernt ist, repräsentieren, worin R²¹ bis R²¹ unabhängig voneinander Wasserstoff oder einen C--C_E-Kohlenwasserstoffrest repräsentieren; oder worin Y -CQ4Q1Q1 repräsentiert, worin Q4, Q5 und Q6 verschieden sind und Q^{ϵ} und Q^{ϵ} wie zuvor Q^{ϵ} , Q^{ϵ} und Q^{ϵ} definiert sind, und wobei Q^{ϵ} (CH_c)_q repräsentiert, von dem eine Bindung mit der Bindung von R4 verknüpft ist, wobei q eine ganze Zahl von 1 bis 4 ist.

- 2. Organische nichtlineare optische Verbindung nach Anspruch 1, worin in der Formel (I) R' Wasserstoff repräsentiert, Z1 Wasserstoff, Rf Rf N-, R7 O-, Rf S-, NC-, Rf OCO-, R10 COO-, O2 N-, R11R10 NOC-, R13 CO-(R*4)N- oder R*5- repräsentiert, Ar eine aromatische Gruppe mit 5 bis 14 Köhlenstoffatomen repräsentiert und B -OH+Amin* repräsentiert, worin Amin* ein optisch aktives Amin. ausgewählt aus der aus 1-Pnenylethylamin, 1-(α-Naphthyl)-ethyl-amin, 1-Phenyl-2-methylethylamin, 1-Phenyl-2-aminopropan und Brucin bestehenden Gruppe, repräsentiert.
- 3. Organische nichtlineare optische Verbindung nach Anspruch 1, worin in der Formel (I) Rt Wasserstoff repräsentiert, Z1 H-, R^ER^EN-, R⁷O-, R⁸S-, NC-, O₂N-, R^eOCO-, R¹⁰COO-, R¹¹R¹²NOCY-, R¹³CO(R¹⁴)Noder R15 repräsentiert, Ar eine aromatische Gruppe mit 6 bis 14 Kohlenstoffatomen und B ein -OH-Amin' repräsentiert, wobei Amin' ein optisch aktives Amin repräsentiert, ausgewahlt aus der aus 2-Amino-1-butanol, 1-Amino-2-propanol, 2-Amino-1-propanol, 2-Amino-1-(p-nitrophenyl)-1,3-propandiol, 2-Dimethyl-amino-1-phenyl-1-benzyl-1-propanol und 1-(N,N-dimethylamino)-1-phenyl-propylamin bestehenden Gruppe.
- Organische nichtlineare optische Verbindung nach Anspruch 1, worin in Formel (I) R1 Wasserstoff repräsentiert, A 25



repräsentiert, worin einer der Reste

Z², Z² und Z⁴ Wasserstoff repräsentiert und die restlichen unabhängig voneinander C;-C-c-Alkyl, R¹6 O-, R¹⁷R¹⁸N-, R¹⁹S oder O₂N- repräsentieren, wobei Ar eine aromatische Gruppe mit 6 bis 14 Kohlenstoffatomen repräsentiert und wobei B einen Rest eines optisch aktiven Amins repräsentiert, welches aus der aus 2-Amino-1-butanol, 1-Amino-2-propanol, 2-Amino-1-propanol, 2-Amino-1-(p-nitrophenyl)-1.3-propandiol, 2-Dimethylamino-1-phenyl-1-benzyl-1-propanol und 1-(N,N-dimethylamino)-1-phenylpropylamin bestehenden Gruppe ausgewählt ist.

5. Organische nichtlineare optische Verbindung nach Anspruch 1, worin in der Formel (I) R1 Wasserstoff repräsentiert. A



repräsentiert, worin einer der Reste

Z², Z² und Z4 Wasserstoff repräsentiert und die restlichen unabhängig voneinander C+-C+o+Alkyl, R'6 O-R¹⁷R¹⁶N-, R¹⁶S- oder O₇N- repräsentieren, worin Ar eine aromatische Gruppe mit 6 bis 14 Köhlenstoffatomen repräsentiert und B einen Rest eines optisch aktiven Amins repräsentiert, ausgewählt aus der aus 1-Phenylethylamin, 1-α-Naphthyl-ethylamin, 1-Phenyl-2-methylethylamin, 1-Phenyl-2aminopropan und Brucin bestehenden Gruppe.

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R°COO-, R°R°NOC-, R°CO(R°4)N- oder R°t repräsentiert, wobei Ar einen aromatischen Ring mit 6 bis 14 Köhlenstoffatomen repräsentiert und wobei BI-NR⁴Y repräsentiert.

- 7. Organische nichtlineare optische Verbindung nach Ansprüch 1, worin in der Formel (I) R* Wasserstoff repräsentiert, A Z*-Ar- repräsentiert, Z* R*R*N- R*O- R*S-, NC-, R*OCO-, R**COO-, R**R*2NOC, R**SCO(R**)N- oder R**E- repräsentiert. Ar eine aromatische Gruppe mit 6 bis 14 Köhlenstoffatomen repräsentiert und worin B -OH+Amin*, wobei Amin* eine optisch aktive α-Aminosäure oder deren Derivate repräsentiert.
- 8. Organische nichtlineare optische Verbindung nach Anspruch 1, worin in der Formel (I) R⁴ Wasserstoff repräsentiert, A

$$z^2$$
 z^3
Ar

repräsentiert, worin einer der Reste

- Z², Z³ und Z⁴ Wasserstoff repräsentieren oder einen substituierten C·-C_E-Alkylrest und wobei die restlichen Reste Z², Z⁵ und Z⁴ zusammen eine Methylen-dioxy-Gruppe repräsentieren, wobei die Dioxy-Gruppen an die benachbarten Positionen an Ar gebunden sind, wobei Ar eine aromatische Gruppe mit 6 bis 14 Kohlenstoffatomen repräsentiert und worin B -OH·Amin* repräsentiert, wobei Amin* ein optisch aktives Amin repräsentiert, ausgewählt aus der aus 1-Phenylethylamin, 1-α-Naphthylethylamin, 1-Phenyl-2-methylethylamin, 1-Phenyl-2-aminopropan, Brucin, 2-Amino-1-butanol, 1-amino-2-propanol, 2-Amino-1-propanol, 2-Amino-1-(p-nitrophenyl)-1,3-propandiol, 2-Dimethyl-amino-1-phenyl-1-benzyl-1-propanol, 1-(N,N-Dimethyl-amino)-1-phenyl-propylamin und einem Rest einer optisch aktiven α-Aminosäure und deren Derivate bestehenden Gruppe; oder -NR⁴ Y.
- 9. Organische nichtlineare optische Verbindung nach Anspruch 1, worin in der Formel (I) R¹ Wasserstoff oder Methyl repräsentiert; A R² repräsentiert, wobei R² Wasserstoff oder eine Alkylgruppe mit 1 bis 12 Kohlenstoffatomen repräsentiert und worin B -OH-Amin¹ repräsentiert, worin Amin¹ ein optisch aktives Amin repräsentiert, ausgewählt aus der aus 1-Phenylethylamin, 1-(α-Naphthyl)ethylamin, 1-Phenyl-2-methylethylamin, 1-Phenyl-2-aminopropan, Brucin; 2-Amino-1-butanol, 1-Amino-2-propanol, 2-Amino-1-propanol, 2-Amino-1-(p-nitrophenyl)-1,3-propandiol, 2-Dimethylamino-1-phenyl-1-benzyl-1-propanol, 1-(N,N-Dimethylamino)-1-phenyl-1-benzyl-1-propanol, 1-(N,N-Dimethylamino)-1-phenyl-propylamin und einem Rest einer optisch aktiven α-Aminosäure und deren Derivate bestehenden Gruppe.
- Organische nichtlineare optische Verbindung nach Anspruch 1, worin in der Formel (I) R' Wasserstoff
 repräsentiert, A

$$(z^5)_r \stackrel{\mathsf{C}}{+} \stackrel{\mathsf{C}}{+} \stackrel{\mathsf{C}}{+}$$

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und B ein -OH-Amin' repräsentiert, wobei Amin' ein optisch aktives Amin repräsentiert ausgewählt aus der aus 1-Phenylethylamin, 1-(α-Naphthyl)ethylamin-1-phenyl-2-methylethylamin, 1-phenyl-2-aminopropan, Brucin, 2-Amino-1-butanol, 1-Amino-2-propanol, 2-Amino-1-propanol, 2-Amino-1-(p-nitrophenyl)-1,3-propandiol, 2-Dimethylamin-1-phenyl-1-benzyl-1-propanol, 1-(N,N-Dimethylamino)-1-phenyl-propylamin und einem Rest einer optisch aktiven α-Aminosäure und deren Derivate bestehenden Gruppe; oder NR⁴Y.

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11. Organische nichtlineare optische Verbindung nach Anspruch 1, worin in der Formel (I) R. Wasserstoff

$$\frac{z^2}{z^3}$$
 Ar

repräsentiert, worin einer der Reste Z^c , Z^d und Z^4 Wasserstoff repräsentiert und die restlichen unabhängig voneinander $C_1 - C_{1/2} - Alkyl$, R^{*c} O_2 , R^{*c} R^{*c} $N_1 - R^{*c}$ $N_2 - R^{*c}$ oder $N_3 - R^{*c}$ $N_3 - R^{*c}$ $N_3 - R^{*c}$ oder $N_3 - R^{*c}$ $N_3 - R^{*c}$ oder $N_3 - R^{*c}$ ode

Revendications

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1. Substance optique non linéaire organique répondant à la formule (I) :

A
$$-(-CR^{1}=CH)\frac{1}{n}$$
 $-CH=C-C-B$ (I)

dans laquelle R¹ représente -H ou -CH₃; n est 0, 1 ou 2; A représente Z¹-Ar-,

$$z^{2}$$
 z^{3}
Ar-, R^{2} - , ou $(z^{5})_{r}$
 C
 C
 C

où Ar représente un groupe aromatique à 6 à 14 chaînons comprenant un cycle hétérocyclique ou un cycle du type bisphénylène ; Z¹ représente H-, R⁵ R⁶ N-, R² O-, R® S-, NC-, R⁵ OCO-, R¹0 COO-, O₂ N-, R¹1R¹²NOC-, R¹3CO(R¹⁴)N- ou R¹⁵- ; Z², Z³, et Z⁴ représentent indépendamment H-, un groupe alkyle en C_1 - C_8 , R¹⁶ O-, R¹²R¹⁶ N-, R¹⁶S-, O_2 N-, ou deux symboles R¹⁶ représentent ensemble R²⁰ CH $\stackrel{\cdot}{\leftarrow}$; R² représente H- ou un groupe alkyle en C_1 - C_1 - C_2 ; R⁵ à R²⁰ représentent indépendamment H-, ou un radical hydrocarbure en C_1 - C_1 - C_2 ; C_2 représente indépendamment H-, un radical hydrocarbure saturé en C_1 - C_2 ; C_2 représente indépendamment H-, un radical hydrocarbure saturé en C_2 - C_3 0 N-, C_3 0 N-, C_4 0 N-, C_5 0 N-, C_5 0 N-, C_6 0 N-,

B représente un groupe -OH+Amine' dans lequel Amine représente une amine optiquement active ; -NR4Y dans lequel R4 représente -H ou une simple liaison ; Y représente -(CH₂)_rCQ¹Q²Q³où p est 0 ou 1 ; Q¹ Q² et Q³ sont différents et représentent -H, un groupe alkyle en C_1 - C_2 , un groupe phényle, naphtyle, -OH, -CH₂OH, -COOR³, -CNR³6-R²7, un radical d'alpha-aminoacide dont un groupe amino est éliminé. R³² à R²¹ représentant indépendamment -H ou un radical hydrocarbure en C_1 - C_2 ; ou Y représente -CQ⁴Q²Q²où Q⁴, Q⁵ et Q⁴ sont différents et Q⁴ et Q³ sont tels que définis pour Q¹, Q⁵ et Q² et Q⁴ et Q⁴ représente (CH₂)_q dont une liaison est liée à la liaison de R⁴ où q est un entier de 1 à 4

2. Substance optique non linéaire organique selon la revendication 1, dans laquelle dans la formule (I), R¹ représente H, Z¹ représente H-, R⁵ R⁶ N-, R² O-, R⁶ S-, NC-, R⁶ OCO-, R˚COO-, O₂ N-, R˚T² NOC-, R˚CO(R˚⁴)N- ou R˚¹-, Ar représente un groupe aromatique en C₅-C-₄ et B représente -OH-Amine¹ où Amine¹ représente une amine optiquement active choisie parmi la 1-phényléthylamine, la 1-(alphanaphtyliéthylamine, la 1-phényl-2-méthyéthylamine, le 1-phényl-2-aminopropane et la brucine

and the control of th

R'3CO(R'4)N- ou R'5-. Ai représente un groupe aromatique en C₁-C-2 et B représente un groupe -OH+Amine* où Amine* représente une amine optiquement active choisie dans le groupe constitué par le 2-aminc-1-putanol, le 1-aminc-2-propanol, le 2-aminc-1-propanol, le 2-aminc-1-(p-nitrophényl)-1,3-propanediol, le 2-diméthyl- amino-1-phényl-1-benzyl-1-propanol et la 1-{N,N-diméthyl-amino}-1-phényl-propylamine

 Sustitance optique non linéaire organique selon la revendication 1, dans laquelle, dans la formule (I), R' représente H, A représente

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$$\begin{array}{c}
z^{2} \\
z^{3} \\
z^{4}
\end{array}$$
 Ar

où un des symboles Z^2 , Z^2 et Z^4 représentent H et les autres représentent indépendamment un groupe alkyle en C_1 - C_1 - C_2 , C_1 - C_2 , C_2 - C_3 - C_4 - C_4 - C_5

 Substance optique non linéaire organique selon la revendication 1, dans laquelle, dans la formule (I) R¹ représente H, A représente



où un des symboles Z^2 , Z^3 et Z^4 représentent H- et les autres représentent indépendamment un groupe alkyle en C_1 - C_{10} , R^{16} O_7 , R^{16} N_7 N_7 ,

- 6. Substance optique non linéaire organique selon la revendication 1, dans laquelle dans la formule (I), R¹ représente H. A représente Z¹Ar-, Z¹ représente H-, R⁵ R6N-, R7O-, R8S, NC-, O₂N-, R9OCO-, R¹OCO-, R¹COCO-, R¹COCO-,
- 7. Substance optique non linéaire organique selon la revendication 1, dans laquelle, dans la formule (I). R' représente H, A représente Z'Ar-, Z' représente R-R'N-,R'O-, R'S-, NC-, R'OCO-, R'OCO-
- Substance optique non linéaire organique selon la revendication 1, dans laquelle, dans la formule (I), R' représente H, A représente



où un des symboles Z^2 , Z^2 et Z^4 représente H ou un groupe alkyle en C_1 - C_2 substitué, les autres représentent ensemble un groupe méthylènedioxy dans lequel les groupes dioxy sont liés aux positions adjacentes de Ar. Ar représente un groupe arbmatique en C_1 - C_2 et B représente un groupe -OH+Amine* dans lequel Amine* représente une amine optiquement active choisie dans le groupe constitué par la 1-phényléthylamine. 1-iaipha-naphtylléthylamine, la 1-phényl-2-méthyléthylamine, le 1-phényl-2-aminopropane, la brucine, le 2-amino-1-butanol, le 1-amino-2-propanol, le 2-amino-1-propanol, le 2-amino-1-phényl-1-3-propanol, le 2-diméthylamino-1-phényl-1-benzyl-1-propanol, la 1-(N,N-diméthylamino)-1-phényl-propylamine et un radical d'un alpha-aminoacide optiquement actif et le dérivé de celui-ci , ou -NR4 Y

9. Substance optique non linéaire organique selon la revendication 1, dans laquelle, dans la formule (I), R' représente H ou CH₃. A représente R², dans lequel R² représente H ou un groupe alkyle en C·-C·₂, et B représente un groupe -OH+Amine' dans leque Amine' représente une amine optiquement active choisie dans le groupe constitué par la 1-(phényl)ethylamine, 1-(alphanaphtyl)éthylamine, la 1-phényl-2-méthyléthylamine, le 1-phényl-2-aminopropane, la brucine ; le 2-amino-1-butanol, le 1-amino-2-propanol, le 2-amino-1-propanol, le 2-amino-1-propanol, le 2-amino-1-phénil-1-benzyl-1-propanol, la 1-(N,N-diméthylamino)-1-phényl-propylamine et un radical d'un alpha-aminoacide optiquement actif et le dérivé de celui-ci.

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Substance optique non linéaire organique selon la revendication 1, dans laquelle, dans la formule (I), R1
représente H, A représente

$$\begin{pmatrix} c - c \\ (z^5)_r & \vdots \\ c & c \\ x \end{pmatrix}$$

- et B représente un groupe -OH+Amine* dans lequel Amine* représente une amine optiquement active choisie dans le groupe constitué par la 1-(phényl)éthylamine, 1-(alphanaphtyl)éthylamine, la 1-phényl-2-méthyléthylamine, le 1-phényl-2-aminopropane, la brucine ; le 2-amino-1-butanol, le 1-amino-2-propanol, le 2-amino-1-propanol, le 2-amino-1-propanol, le 2-amino-1-phényl-1-benzyl-1-propanol, la 1-(N,N-diméthylamino)-1-phényl-propylamine et un radical d'un alpha-aminoacide optiquement actif et les dérivés de celui-ci ; ou NR⁴ Y.
- Substance optique non linéaire organique selon la revendication 1, dans laquelle, dans la formule (I), R¹
 représente H, A représente

$$z^2$$
, $z^3 \rightarrow Ar$

où un des symboles Z², Z³ et Z⁴ représentent H- et les autres représentent indépendamment un groupe alliyle en C--C-c R² R² N-, R² N-, R² S, où C₅ N-, Ar représente un groupe aromatique en C₆-C-c et B représente un groupe - OH+Amine* dans lequel Amine* représente des alpha-aminoacides optiquement actifs et leurs dérivés.